PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference 040826woMemh	FOR FURTHER ACTION See Form PCT/PEA/416						
International application No. PCT/EP2004/003115	International filing date (da 24.03.2004	nternational filing date (day/month/year) Priority date (24.03.2004 Priority date (24.03.2003					
International Patent Classification (IPC) or national classification and IPC A61K38/19							
Applicant IPF PHARMACEUTICALS GMBH et al.							
 This report is the international preliminary examination report, established by this International Preliminary Examining Authority under Article 35 and transmitted to the applicant according to Article 36. 							
2. This REPORT consists of a total of							
3. This report is also accompanied b	y ANNEXES, comprising	: 	- a fallerine				
a. 🛛 sent to the applicant and to	a. Sent to the applicant and to the International Bureau) a total of 2 sheets, as follows:						
sheets of the description, claims and/or drawings which have been amended and are the basis of this report and/or sheets containing rectifications authorized by this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions).							
sheets which supersed beyond the disclosure Supplemental Box.	sheets which supersede earlier sheets, but which this Authority considers contain an amendment that goes heyond the disclosure in the international application as filed, as indicated in item 4 of Box No. I and the						
b. (sent to the International E	— u						
Box Helating to Sequence	Listing (See Seetier SE						
4. This report contains indications re	elating to the following ite	ms:					
☐ Box No. I Basis of the opi	inion						
☐ Box No. II Priority							
		d to novelty, inventive :	step and industrial applicability				
☐ Box No. IV Lack of unity of	invention		to continue atom or industrial				
applicability; cit	applicability; citations and explanations supporting such statement						
☐ Box No. VI Certain docume		4:					
	in the international appli						
☐ Box No. VIII Certain observe	☐ Box No. VIII Certain observations on the international application						
Date of submission of the demand		Date of completion of thi	s report				
Date of Submission of the Communication							
24.01.2005		15.04.2005					
Name and mailing address of the international preliminary examining authority:		Authorized Officer	Josephiches Polonica.				
European Patent Office		Didelon, F					
Tel. +49 89 2399 - 0 Tx: 523 Fax: +49 89 2399 - 4465	656 epmu d	Telephone No. +49 89 2	399-7332				

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/003115

	Box	No. I Basis of the report						
1.	With	th regard to the language , this report is based on the international application in the language in which it d, unless otherwise indicated under this item.						
		This report is based on translations from the original language into the following language, which is the language of a translation furnished for the purposes of: international search (under Rules 12.3 and 23.1(b)) publication of the international application (under Rule 12.4)						
		international preliminary examination (under Rules 55.2 and/or 55.3)						
2.	ha	h regard to the elements* of the international application, this report is based on <i>(replacement sheets which</i> The been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this Ort as "originally filed" and are not annexed to this report):						
	Des	scription, Pages						
	1-12	2 as originally filed						
	Cla	ims, Numbers						
	1-9	filed with telefax on 24.01.2005						
	Dra	awings, Sheets						
	1/1	as originally filed						
		a sequence listing and/or any related table(s) - see Supplemental Box Relating to Sequence Listing						
3.		The amendments have resulted in the cancellation of: ☐ the description, pages ☐ the claims, Nos. ☐ the drawings, sheets/figs ☐ the sequence listing (specify): ☐ any table(s) related to sequence listing (specify):						
4.	ha Su	d not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the applemental Box (Rule 70.2(c)). the description, pages the claims, Nos. the drawings, sheets/figs the sequence listing (specify): any table(s) related to sequence listing (specify):						
	*	If item 4 applies, some or all of these sheets may be marked "superseded."						

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/003115

	арр	licability		nion with regard to novelty, inventive step and industrial		
 The questions whether the claimed invention appears to be novel, to involve an inventive obvious), or to be industrially applicable have not been examined in respect of: 				tion appears to be novel, to involve an inventive step (to be non- ave not been examined in respect of:		
		the entire international application	entire international application,			
	\boxtimes	claims Nos. 6-9 (IA)				
		because:				
	×	the said international application, or the said claims Nos. 6-9 relate to the following subject matter which does not require an international preliminary examination (specify):				
		see separate sheet				
		that no meaningful opinion coul	on, claims or drawings <i>(indicate particular elements below)</i> or said claims Nos. are so unclear ningful opinion could be formed <i>(specify)</i> :			
		the claims, or said claims Nos. could be formed.	s Nos. are so inadequately supported by the description that no meaningful opinion			
		no international search report h	has been established for the said claims Nos.			
		the nucleotide and/or amino aci C of the Administrative Instruct	mino acid sequence listing does not comply with the standard provided for in Annex			
		the written form		has not been furnished		
				does not comply with the standard		
		the computer readable form		has not been furnished		
		·		does not comply with the standard		
		the tables related to the nucleon not comply with the technical r	I to the nucleotide and <i>l</i> or amino acid sequence listing, if in computer readable form or he technical requirements provided for in Annex C- <i>bis</i> of the Administrative Instructio			
		See separate sheet for further	detai	ils		

Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N) Yes: Claims 1-9

No: Claims

Inventive step (IS) Yes: Claims 1-9

No: Claims

Industrial applicability (IA) Yes: Claims 1-5

No: Claims

2. Citations and explanations (Rule 70.7):

see separate sheet

Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

Claims 6-9 relate to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Said claim seem to depend on use claims but are formulated as method claims. Consequently, no opinion will be formulated with respect to the industrial applicability of the subjectmatter of these claims (Article 34(4)(a)(I) PCT).

Re Item V

Reasoned statement under Rule 66.2(a)(ii) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- Reference is made to the following documents: 1.
 - D1: MOLLER C ET AL: "Expression and function of chemokine receptors in human multiple myeloma." LEUKEMIA (BASINGSTOKE), vol. 17, no. 1, 20 January 2003 (2003-01-20), pages 203-210, XP002252455 ISSN: 0887-6924
 - D2: WRIGHT DOUGLAS E ET AL: "Hematopoietic stem cells are uniquely selective in their migratory response to chemokines." JOURNAL OF EXPERIMENTAL MEDICINE, vol. 195, no. 9, 6 May 2002 (2002-05-06), pages 1145-1154, XP002252456 May 6, 2002 ISSN: 0022-1007
 - D3: WO 00/46248 A (SCHERING CORP) 10 August 2000 (2000-08-10)
 - D4: WO 01/17558 A (SCHERING CORP) 15 March 2001 (2001-03-15)
 - D5: US 2003/049696 A1 (GEORGE THADDEUS C ET AL) 13 March 2003 (2003-03-13)
 - D6: WO 00/73432 A (CORNELL RES FOUNDATION INC) 7 December 2000 (2000-12-07)
 - D7: WO 00/03016 A (CONNEX GMBH ;REITER CHRISTIAN (DE)) 20 January 2000 (2000-01-20)
 - D8: CA 2 256 250 A (UAB FOUNDATION, US) 17 June 1999 (1999-06-17)

Unless otherwise indicated, the relevant passages in the cited documents are the ones

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indicated in the Search Report.

Novelty and inventive step :

D1 reveals that multiple myelomas cells migrate in response to chemokines SDF-1, MIP-1alpha and RANTES. This could be used as a new treatment for multiple myeloma. However it does not disclose stem cells.

D2 reports that, contrary to the present application, hematopoietic stem cells only respond to SDF-1, ligand of CXCR-4 receptor but not to other chemokines. However the combination of both SDF-1 and other chemokines is effective. The specific chemokine ligands of CCR-3 -6 an d-8 are however not effective on the migration of hematopoietic stem cells.

In D3 and D4, MIP-3alpha or CCR6 agonist are contemplated for use in some diseases, including skin grafts. However stem cells are not encompassed in the disclosure and the goal is not of enhancing homing of stem cells but to reduce inflammatory response against said grafts.

D5 discloses the trafficking of regulatory T-cells is modulated by agonists of CCR6, e.g., MIP-3alpha or LARC. This finds usefulness in transplantation and in modulation of immune response. This document however does not deal with the migration of stem cells.

D6 describes SDF-1 and MIP-3alpha, delivered through adenoviral vectors, and able to attract dendritic cells in vitro and are thus used to enhance immunity, in particular to combat cancers.

Again stem cells are not encompassed in said document.

D7 reveals that RANTES, MIP-1alpha (CCR3 agonists), MIP-1beta (CCR8 agonist) are used as pharmaceutical compositions for prevention of autoimmune diseases, immune deficiencies, infectious diseases, suppression of immune response in graft rejection. It does not relate however to the improvement of stem cell migration for increasing transplantation efficiency.

D8 disclosed chemokines such as RANTES, MIP-lalpha, MIP-lbeta used as mucosal immune enhancers.

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

International application No.

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It is understood that a novel cellular mechanism has been discovered, in that the ligands of different chemokines receptors have been found to potentiate the effects of the CXCR4 receptor ligand SDF-1 for improving stem cell transplantation.

Claim 1 which satisfactorily defines the medical application underlying the present application "treatment of progenitor/stem cells prior to and/or in the course of transplantation" is thus considered as novel and inventive because the prior art does not reveal nor suggest the treatment of progenitor or stem cells used in the application for improving transplantation.

- 3. For the assessment of the present claims 6-9 on the question whether they are industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.
- 4. For the sake of completeness, it must be made clear that in front of some regional Authorities, human embryonic stem cells (see claim 3) are not to be encompassed within the scope of the present application.

S. 9/20NR. 8775

JC14 Rec'd PCT/PYC 23 SEP 2005

Claims

The use of at least one agonist of receptors selected from the group 1. consisting of the CCR3, CCR6 or CCR8 receptor or combinations thereof and a pharmaceutically acceptable carrier for treatment of progenitor and stem cells prior to and/or in the course of transplantation of the cells wherein the agonist is selected from the group consisting

of receptor CCR3: Eotaxin; Eotaxin-2; Eotaxin-3; Hemofiltrate CC-Chemokine-1 (HCC-1); Hemofiltrate CC Chemokine-2 Macrophage Inflammatory Protein - 10 (MIP-10); Regulated on Activation Normally T-Cell Express and Secreted (RANTES); Monocyte Chemoattractant Protein - 2 (MCP-2); Monocyte Chemoattractant Protein - 3 (MCP-3); Monocyte Chemoattractant Protein - 4 (MCP-4); 2-[(6-amino-2-benzothiazolyl)thio]-N-[1-[(3,4-dichlorylphenyl)methyl]-4piperidinyl] acetamide;

of receptor CCR6: Macrophage Inflammatory Protein - 3a (MIP-3a);

of receptor CCR8: I309; Macrophage Inflammatory Protein - 1ß (MIP-1β); LAG-1; Thymus and Activation Regulated Chemokine (TARC); viral Macrophage Inflammatory Protein - I (vMIP-I); as well as derivatives therof keeping their agonist abilities.

- The use of claim 1 for improving the homing of stem cells. 2.
- The use according to one or more of the foregoing claims for the 3. transplantation of hematopoietic progenitor and stem cells, umbilical cord blood and placental stem and progenitor cells, liver stem and progenitor cells (oval cells), mesenchymal stem and progenitor cells, endothelial progenitor cells, skeletal muscle stem and progenitor cells (satellite cells), smooth muscle stem and progenitor cells, intestinal stem and progenitor cells, embryonic stem cells, and genetically modified embryonic stem cells, adult islet/beta stem- and progenitor cell, epidermal progenitor and stem cells, keratinocyte stem cells of

- 2 -

cornea, skin and hair follicles, olfactory (bulb) stem and progenitor cells and side population cells from diverse adult tissues.

- The use according one or more of the foregoing claims to increase the sensitivity of hematopoietic stem cells to SDF-1 induced cellular signals.
- 5. The use according one or more of the foregoing claims for the treatment of leukemias, lymphoproliferative disorders, aplastic anemia, congenital disorders of the bone marrow, solid tumors, autoimmune disorders, inflammatory diseases, primary immunodeficiencies, primary systemic amyloidosis, systemic sclerosis, heart diseases, liver diseases, neurodegenerative diseases, multiple sclerosis, M. Parkinson, stroke, spinal cord injury diabetes mellitus, bone diseases, skin diseases, replacement therapy of the skin, retina or cornea, other congenital disorders, vessel diseases like atherosclerosis or cardiovascular disease.
- The method of the foregoing claim wherein the host patient are not conditioned.
- The method of claim 6 wherein the host patient is conditioned under sublethal, lethal, or supralethal conditions.
- The method according to claim 7 wherein sublethal, lethal, or supralethal conditions include treatment with total body irradiation, optionally followed by treatment with myeloablative or immunosuppressive agents.
- The method according to any one of the claims 7 or 8 wherein sublethal, lethal, or supralethal conditions include myeloablative or immunosuppressive treatment without total body irradiation.